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(71) Applicant: NOVO NORDISK BIOTECH, INC. [US/US]; 1445 Drew Avenue, Davis, CA 95616-4880 (US).

(72) Inventors: BERKA, Randy, M.; 3609 Modoc Place, Davis, CA 95616 (US). XU, Feng; 1534 Carmel Valley Drive, Woodland, CA 95776 (US). THOMPSON, Sheryl, A.; 5207 Colwell, Davis, CA 95616 (US).

(74) Agents: ZELSON, Steve, T. et al.; Novo Nordisk of North America, Inc., Suite 6400, 405 Lexington Avenue, New York, NY 10174 (US). (81) Designated States: AM, AU, BB, BG, BR, BY, CA, CN, CZ, EE, FI, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LT, LV, MD, MG, MN, MX, NO, NZ, PL, RO, RU, SD, SG, SI, SK, TJ, TM, TT, UA, UG, UZ, VN, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG), ARIPO patent (KE, MW, SD, SZ, UG).

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(54) Title: PURIFIED SCYTALIDIUM LACCASES AND NUCLEIC ACIDS ENCODING SAME

(57) Abstract

The present invention relates to isolated nucleic acid constructs containing a sequence encoding a Scytalidium laccase, and the laccase proteins encoded thereby.

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PURIFIED SCYTALIDIUM LACCASES AND NUCLEIC ACIDS ENCODING SAME

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Field of the Invention

The present invention relates to isolated nucleic acid fragments encoding a fungal oxidoreductase enzyme and the purified enzymes produced thereby. More particularly, the invention relates to nucleic acid fragments encoding a phenol oxidase, specifically a laccase, of a thermophilic fungus, Scytalidium.

15 Background of the Invention

Laccases (benzenediol:oxygen oxidoreductases) are multi-copper containing enzymes that catalyze the oxidation of phenolics. Laccase-mediated oxidations result in the production of aryloxy-radical intermediates from suitable 20 phenolic substrate; the ultimate coupling of the intermediates so produced provides a combination of dimeric, oligomeric, and polymeric reaction products. Such reactions are important in nature in biosynthetic pathways which lead to the formation of melanin, alkaloids, toxins, lignins, and 25 humic acids. Laccases are produced by a wide variety of fungi, including ascomycetes such as Aspergillus, Neurospora, and Podospora, the deuteromycete Botrytis, and basidiomycetes such as Collybia, Fomes, Lentinus, Pleurotus, Trametes, and perfect forms of Rhizoctonia. Laccase 30 exhibits a wide range of substrate specificity, and each different fungal laccase usually differs only quantitatively from others in its ability to oxidize phenolic substrates. Because of the substrate diversity, laccases generally have found many potential industrial applications. Among these

are lignin modification, paper strengthening, dye transfer inhibition in detergents, phenol polymerization, juice manufacture, phenol resin production, and waste water treatment.

Although the catalytic capabilities are similar, laccases made by different fungal species do have different temperature and pH optima, and these may also differ depending on the specific substrate. A number of these fungal laccases have been isolated, and the genes for 10 several of these have been cloned. For example, Choi et al. (Mol. Plant-Microbe Interactions 5: 119-128, 1992) describe the molecular characterization and cloning of the gene encoding the laccase of the chestnut blight fungus, Cryphonectria parasitica. Kojima et al. (J. Biol. Chem. 15 <u>265</u>: 15224-15230, 1990; JP 2-238885) provide a description of two allelic forms of the laccase of the white-rot basidiomycete Coriolus hirsutus. Germann and Lerch (Experientia 41: 801,1985; PNAS USA 83: 8854-8858, 1986) have reported the cloning and partial sequencing of the 20 Neurospora crassa laccase gene. Saloheimo et al.(J. Gen. Microbiol. 137: 1537-1544, 1985; WO 92/01046) have disclosed a structural analysis of the laccase gene from the fungus Phlebia radiata.

Attempts to express laccase genes in heterologous

fungal systems frequently give very low yields (Kojima et al., supra; Saloheimo et al., Bio/Technol. 2: 987-990,

1991). For example, heterologous expression of Phlebia radiata laccase in Trichoderma reesei gave only 20 mg per liter of active enzyme (Saloheimo, 1991, supra). Although

laccases have great commercial potential, the ability to express the enzyme in significant quantities is critical to their commercial utility. At the present time there are no laccases which are expressed at high levels in commercially utilized hosts such as Aspergillus. Thus, the need exists

for a laccase which can be produced in commercially useful (i.e., gram per liter or more) quantities. The present invention fulfills such a need.

5 Summary of the Invention

The present invention relates to a DNA construct containing a nucleic acid sequence encoding a Scytalidium laccase. The invention also relates to an isolated laccase encoded by the nucleic acid sequence. Preferably, the

10 laccase is substantially pure. By "substantially pure" is meant a laccase which is essentially (i.e.,≥90%) free of other non-laccase proteins.

In order to facilitate production of the novel laccase, the invention also provides vectors and host cells

comprising the claimed nucleic acid fragment, which vectors and host cells are useful in recombinant production of the laccase. The nucleic acid fragment is operably linked to transcription and translation signals capable of directing expression of the laccase protein in the host cell of

choice. A preferred host cell is a fungal cell, most preferably of the genus Aspergillus. Recombinant production of the laccase of the invention is achieved by culturing a host cell transformed or transfected with the nucleic acid fragment of the invention, or progeny thereof, under

conditions suitable for expression of the laccase protein, and recovering the laccase protein from the culture.

The laccases of the present invention are useful in a number of industrial processes in which oxidation of phenolics is required. These processes include lignin manipulation, juice manufacture, phenol polymerization and phenol resin production.

Brief Description of the Figures

Figure 1 illustrates the nucleotide (SEQ ID NO: 1) and amino acid (SEQ ID NO: 2) sequence of Scytalidium thermophila laccase. Letters without corresponding amino acids in the nucleotide sequence indicate the position of introns.

Figure 2 illustrates the construction of plasmid pShTh15.

Figure 3 illustrates the restriction map of a XhoI insert in pShTh6 which contains the S. thermophilum

10 laccase(lccS) gene. The approximate position of the lccS coding region is indicated by a solid black line.

Figure 4 illustrates the pH profiles of the laccase activity with syringaldazine(squares) and 2,2" azinobis(3-ethylbenzothiazoline-6-sulfonic acid)(circles) as substrate.

15 Figure 5 illustrates the thermostability in B&R buffers of the laccase at pH 2.7, 6.1, and 9.0. Preincubation times are 1 hour. Activities are assayed by ABTS oxidation at 20°C in B&R buffer, pH 4.1.

20 Detailed Description of the Invention

Scytalidium thermophilum is a thermophilic deuteromycete, and a member of the Torula-Humicola complex which are recognized as dominant species in mushroom compost. Other members of the complex include Humicola grisea Traaen var. thermoidea Cooney & Emerson, H. insolens Cooney & Emerson, and Torula thermophila Cooney & Emerson, the latter of which has been reassigned to Scytalidium thermophilum by Austwick (N.Z. J. Agric. Res. 19: 25-33, 1976). Straatsma and Samson (Mycol. Res. 97: 321-328, 1993) have recently determined that both H. grisea var.thermoides and H. insolens should be considered as examples of the species Scytalidium thermophilum as well. S. indonesiacum (Hedger et al., Trans. Brit Mycol.Soc. 78: 366-366, 1982) may also be synonymous with S. thermophilum. Members of the

complex are known to be producers of thermostable cellulase and ß-glucosidase enzymes(Rao and Murthy, Ind. J. Biochem. Biophys. 25: 687-694, 1988; Hayashida and Yoshioka, Agric. Biol. Chem. 44: 1721-1728, 1980). However, there have been no previous reports of the production of a laccase by Scytalidium, or any of the noted synonymous species. It has now been determined that not only does Scytalidium produce a laccase, but the gene encoding this laccase can be used to produce large yields of the enzyme in convenient host systems such as Aspergillus.

To identify the presence of a laccase gene in Scytalidium, a 5' portion of the Neurospora crassa laccase gene(lcc1) is used as a probe, under conditions of mild stringency, in southern hybridization of total genomic DNA 15 of different fungal species. An approximately 3 kb laccase specific sequence is detected in the Scytalidium DNA. N. crassa fragment is then used to screen about 12,000 plaques of an s. thermophilum genomic DNA library in a λ EMBL4 bacteriophage cloning vector. Nine plaques strongly 20 hybridize with the probe; from these nine, DNA is isolated from four. Each of these clones contains a 3kb BamHI fragment corresponding to the one initially identified in the southern blot of genomic DNA. One of the fragments is subcloned into a pBluescript vector; however, DNA sequencing 25 shows only a portion of the gene to be on this fragment. A 6kb fragment XhoI fragment from the same phage contains the whole Iccs gene, and this is then subcloned into pBluescript to derive plasmid pShTh6. A restriction map of the 6 kb insert is shown in Figure 3.

Once the sequence is determined, the positions of introns and exons within the gene is assigned based on alignment of the deduced amino acid sequence to the corresponding N. crassa laccase gene product. From this comparison, it appears that the gene (lccs) of S.

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thermophilum is composed of seven exons(243, 91, 70, 1054 and 390 nucleotides) punctuated by four small introns (63, 58, 55 and 65 nucleotides). The coding region, excluding intervening sequences is very GC-rich(60.8% G+C) and encodes a preproenzyme of 616 amino acids: a 21 amino acid signal peptide and a 24 amino acid propeptide. The sequence of the S. thermophilum gene and the predicted amino acid sequence is shown in Figure 1 (SEQ ID NOS: 1 and 2)

The laccase gene is then used to create an expression

vector for transformation of Aspergillus host cells. The

vector, pShTh15 contains the A. oryzae TAKA-amylase promoter

and the A. niger glaA terminator regions. The construction

of pShTh15 is outlined in Figure 2. Aspergillus cells are

cotransformed with the expression vector and a plasmid

containing the pyrG or amdS selectable marker.

Transformants are selected on the appropriate selective

medium containing ABTS. Laccase-producing colonies exhibit a

green halo and are readily isolatable. Selected

transformants are grown up in shake flasks and culture

broths tested for laccase activity by the syringaldazine

method. Shake flask cultures are capable of producing 50 or

more mg/liter of laccase, and in fermentors, yields of over

1.6 g/liter are observed.

According to the invention, a Scytalidium gene
25 encoding a laccase can be obtained by methods described
above, or any alternative methods known in the art, using
the information provided herein. The gene can be expressed,
in active form, using an expression vector. A useful
expression vector contains an element that permits stable
30 integration of the vector into the host cell genome or
autonomous replication of the vector in a host cell
independent of the genome of the host cell, and preferably
one or more phenotypic markers which permit easy selection
of transformed host cells. The expression vector may also

include control sequences encoding a promoter, ribosome binding site, translation initiation signal, and, optionally, a repressor gene or various activator genes. To permit the secretion of the expressed protein, nucleotides 5 encoding a signal sequence may be inserted prior to the coding sequence of the gene. For expression under the direction of control sequences, a laccase gene to be used according to the invention is operably linked to the control sequences in the proper reading frame. Promoter 10 sequences that can be incorporated into plasmid vectors, and which can direct the transcription of the laccase gene, include but are not limited to the prokaryotic ß-lactamase promoter (Villa-Kamaroff, et al., 1978, Proc. Natl. Acad. Sci. U.S.A. 75:3727-3731) and the tac promoter (DeBoer, et 15 al., 1983, Proc. Natl. Acad. Sci. U.S.A. <u>80</u>:21-25). Further references can also be found in "Useful proteins from recombinant bacteria" in Scientific American, 1980, 242:74-94; and in Sambrook et al., Molecular Cloning, 1989.

The expression vector carrying the DNA construct of the
invention may be any vector which may conveniently be
subjected to recombinant DNA procedures, and the choice of
vector will typically depend on the host cell into which it
is to be introduced. Thus, the vector may be an autonomously
replicating vector, i.e. a vector which exists as an
extrachromosomal entity, the replication of which is
independent of chromosomal replication, e.g. a plasmid, or
an extrachromosomal element, minichromosome or an artificial
chromosome. Alternatively, the vector may be one which, when
introduced into a host cell, is integrated into the host
cell genome and replicated together with the chromosome(s)
into which it has been integrated.

In the vector, the DNA sequence should be operably connected to a suitable promoter sequence. The promoter may be any DNA sequence which shows transcriptional activity in

the host cell of choice and may be derived from genes encoding proteins either homologous or heterologous to the host cell. Examples of suitable promoters for directing the transcription of the DNA construct of the invention, 5 especially in a bacterial host, are the promoter of the lac operon of E.coli, the Streptomyces coelicolor agarase gene dagA promoters, the promoters of the Bacillus licheniformis α-amylase gene (amyL), the promoters of the Bacillus stearothermophilus maltogenic amylase gene (amyM), the 10 promoters of the Bacillus amyloliquefaciens α-amylase (amyQ), or the promoters of the Bacillus subtilis xylA and xylB genes. In a yeast host, a useful promoter is the eno-1 promoter. For transcription in a fungal host, examples of useful promoters are those derived from the gene encoding A. 15 oryzae TAKA amylase, Rhizomucor miehei aspartic proteinase, A. niger neutral α -amylase, A. niger acid stable α -amylase, A. niger or A. awamori glucoamylase (glaA), Rhizomucor miehei lipase, A. oryzae alkaline protease, A. oryzae triose phosphate isomerase or A. nidulans acetamidase. Preferred 20 are the TAKA-amylase and glaA promoters.

The expression vector of the invention may also comprise a suitable transcription terminator and, in eukaryotes, polyadenylation sequences operably connected to the DNA sequence encoding the laccase of the invention. Termination and polyadenylation sequences may suitably be derived from the same sources as the promoter. The vector may further comprise a DNA sequence enabling the vector to replicate in the host cell in question. Examples of such sequences are the origins of replication of plasmids pUC19, pACYC177, pUB110, pE194, pAMB1 and pIJ702.

The vector may also comprise a selectable marker, e.g. a gene the product of which complements a defect in the host cell, such as the dal genes from B.subtilis or B.li-cheniformis, or one which confers antibiotic resistance such as ampicillin, kanamycin, chloramphenicol or tetracycline resistance. Examples of Aspergillus selection markers include amdS, pyrG, argB, niaD, sC, and hygB a marker giving rise to hygromycin resistance. Preferred for use in an Aspergillus host cell are the amdS and pyrG markers of A. nidulans or A. oryzae. A frequently used mammalian marker is the dihydrofolate reductase (DHFR) gene. Furthermore, selection may be accomplished by co-transformation, e.g. as described in WO 91/17243.

15 It is generally preferred that the expression gives rise to a product that is extracellular. The laccases of the present invention may thus comprise a preregion permitting secretion of the expressed protein into the culture medium. If desirable, this preregion may be native to the laccase of 20 the invention or substituted with a different preregion or signal sequence, conveniently accomplished by substitution of the DNA sequences encoding the respective preregions. For example, the preregion may be derived from a glucoamylase or an amylase gene from an Aspergillus species, 25 an amylase gene from a Bacillus species, a lipase or proteinase gene from Rhizomucor miehei, the gene for the α factor from Saccharomyces cerevisiae or the calf preprochymosin gene. Particularly preferred, when the host is a fungal cell, is the preregion for A. oryzae TAKA 30 amylase, A. niger neutral amylase, the maltogenic amylase form Bacillus NCIB 11837, B. stearothermophilus α-amylase, or Bacillus licheniformis subtilisin. An effective signal sequence is the A. oryzae TAKA amylase signal, the

Rhizomucor miehei aspartic proteinase signal and the Rhizomucor miehei lipase signal.

The procedures used to ligate the DNA construct of the invention, the promoter, terminator and other elements,

5 respectively, and to insert them into suitable vectors containing the information necessary for replication, are well known to persons skilled in the art (cf., for instance, Sambrook et al. Molecular Cloning, 1989).

The cell of the invention either comprising a DNA construct or an expression vector of the invention as defined above is advantageously used as a host cell in the recombinant production of a enzyme of the invention. The cell may be transformed with the DNA construct of the invention, conveniently by integrating the DNA construct in the host chromosome. This integration is generally considered to be an advantage as the DNA sequence is more likely to be stably maintained in the cell. Integration of the DNA constructs into the host chromosome may be performed according to conventional methods, e.g. by homologous or heterologous recombination. Alternatively, the cell may be transformed with an expression vector as described above in connection with the different types of host cells.

The host cell may be selected from prokaryotic cells, such as bacterial cells. Examples of suitable bacteria are gram positive bacteria such as Bacillus subtilis, Bacillus licheniformis, Bacillus lentus, Bacillus brevis, Bacillus stearothermophilus, Bacillus alkalophilus, Bacillus amyloliquefaciens, Bacillus coagulans, Bacillus circulans, Bacillus lautus, Bacillus megaterium, Bacillus thuringiensis, or Streptomyces lividans or Streptomyces murinus, or gram negative bacteria such as E.coli. The transformation of the bacteria may for instance be effected

by protoplast transformation or by using competent cells in a manner known per se.

The host cell may also be a eukaryote, such as mammalian cells, insect cells, plant cells or preferably 5 fungal cells, including yeast and filamentous fungi. For example, useful mammalian cells include CHO or COS cells. yeast host cell may be selected from a species of Saccharomyces or Schizosaccharomyces, e.g. Saccharomyces cerevisiae. Useful filamentous fungi may selected from a 10 species of Aspergillus, e.g. Aspergillus oryzae or Aspergillus niger. Alternatively, a strain of a Fusarium species, e.g. F. oxysporum, can be used as a host cell. Fungal cells may be transformed by a process involving protoplast formation and transformation of the protoplasts fol-15 lowed by regeneration of the cell wall in a manner known per se. A suitable procedure for transformation of Aspergillus host cells is described in EP 238 023. A suitable method of transforming Fusarium species is described by Malardier et al., 1989.

The present invention thus provides a method of producing a recombinant laccase of the invention, which method comprises cultivating a host cell as described above under conditions conducive to the production of the enzyme and recovering the enzyme from the cells and/or culture

25 medium. The medium used to cultivate the cells may be any conventional medium suitable for growing the host cell in question and obtaining expression of the laccase of the invention. Suitable media are available from commercial suppliers or may be prepared according to published formulae

30 (e.g. in catalogues of the American Type Culture Collection).

The resulting enzyme may be recovered from the medium by conventional procedures including separating the cells from the medium by centrifugation or filtration, precipitat-

PCT/US95/06816 WO 95/33837

ing the proteinaceous components of the supernatant or filtrate by means of a salt, e.g. ammonium sulphate, followed by purification by a variety of chromatographic procedures, e.g. ion exchange chromatography, gel filtration 5 chromatography, affinity chromatography, or the like. Preferably, the isolated protein is about 90% pure as determined by SDS-PAGE, purity being most important in food, juice or detergent applications.

In a particularly preferred embodiment, the expression 10 of laccase is achieved in a fungal host cell, such as Aspergillus. As described in detail in the following examples, the laccase gene is ligated into a plasmid containing the Aspergillus oryzae TAKA α -amylase promoter, and the Aspergillus nidulans amdS selectable marker. 15 Alternatively, the amdS may be on a separate plasmid and used in co-transformation. The plasmid (or plasmids) is used to transform an Aspergillus species host cell, such as A. oryzae or A. niger in accordance with methods described

in Yelton et al. (PNAS USA 81: 1470-1474,1984).

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Those skilled in the art will recognize that the invention is not limited to use of the nucleic acid fragments specifically disclosed herein, for example, in Figure 1. It will also be apparent that the invention encompasses those nucleotide sequences that encode the same 25 amino acid sequences as depicted in Figure 1, but which differ from those specifically depicted nucleotide sequences by virtue of the degeneracy of the genetic code. Also, reference to Figure 1, in the specification and the claims will be understood to encompass both the genomic sequence 30 depicted therein as well as the corresponding cDNA and RNA sequences, and the phrases "DNA construct" and "nucleic acid sequences" as used herein will be understood to encompass all such variations. "DNA construct" shall generally be understood to mean a DNA molecule, either single- or doublestranded, which may be isolated in partial form from a naturally occurring gene or which has been modified to contain segments of DNA which are combined and juxtaposed in a manner which would not otherwise exist in nature.

In addition, the invention also encompasses other Scytalidium laccases, including alternate forms of laccase which may be found in S. thermophilum and as well as laccases which may be found in other fungi which are synonyms or fall within the definition of Scytalidium 10 thermophilum as defined by Straatsma and Samson, 1993, These include S. indonesiacum, Torula thermophila, Humicola brevis var. thermoidea, Humicola brevispora, H. grisea var. thermoidea, Humicola insolens, and Humicola lanuginosa (also known as Thermomyces lanuginosus). 15 invention also provides the means for isolation of laccase genes from other species of Scytalidium, such as S. acidophilum, S. album, S. aurantiacum, S. circinatum, S. flaveobrunneum, S. hyalinum, S. lignicola, and S. uredinicolum. Identification and isolation of laccase genes 20 from sources other than those specifically exemplified herein can be achieved by utilization of the methodology described in the present examples, with publicly available Scytalidium strains. Alternately, the sequence disclosed herein can be used to design primers and/or probes useful in 25 isolating laccase genes by standard PCR or southern hybridization techniques, using the same publicly available strains. Examples of such publicly available strains include, from the American Type Culture Collection, ATCC 16463, 28085, 36346, 48409, 66938(S. thermophilum);24569(S. 30 acidophilum); 16675(S. album); 22477(S. aurantiacum); 66463(S. circinatum);13212(S. flavo-brunneum); 52297(S. fulvum); 38906(S. hyalinum); 46858(S. indonesiacum); 18984(S. indonesiacum); 32382(S. uredinaolum); from the International Mycological Institute (IMI; United Kingdom),

IMI 243 118(S. thermophilum); from Centraalbureau voor Schimmelcultures (CBS; Netherlands) CBS 183.81, 671.88(S. thermophilum) 367.72(S. acidophilum); 372.65(S. album);374.65(S. aurantiacum);654.89(S. circinatum);244.59 5 (S. flavo-brunneum);145.78 (S. hyalinum);259.81(S. indonesiacum);233.57(S. lignicola);171.40(S. terminale); 616.84(S. muscorum); from Deutsche Sammlung von Mikroorganismenn und Zellkulturen (DSM; Germany) DSM 2842(S thermophilum); DSM 2695 (S. lignicola). The invention also 10 encompasses any variant nucleotide sequence, and the protein encoded thereby, which protein retains at least about an 80%, preferably about 85%, and most preferably at least about 90-95% homology with the amino acid sequence depicted in Figure 1, and which qualitatively retains the laccase 15 activity of the sequence described herein. Useful variants within the categories defined above include, for example, ones in which conservative amino acid substitutions have been made, which substitutions do not significantly affect the activity of the protein. By conservative substitution is 20 meant that amino acids of the same class may be substituted by any other of that class. For example, the nonpolar aliphatic residues Ala, Val, Leu, and Ile may be interchanged, as may be the basic residues Lys and Arg, or the acidic residues Asp and Glu. Similarly, Ser and Thr are 25 conservative substitutions for each other, as are Asn and It will be apparent to the skilled artisan that such substitutions can be made outside the regions critical to the function of the molecule and still result in an active enzyme. Retention of the desired activity can readily be 30 determined by conducting a standard ABTS oxidation method, such as is described in the present examples.

The protein can be used in number of different industrial processes. These processes include polymerization of lignin, both Kraft and lignosulfates, in

solution, in order to produce a lignin with a higher molecular weight. A neutral/alkaline laccase is a particular advantage in that Kraft lignin is more soluble at higher pHs. Such methods are described in, for example, Jin et al., Holzforschung 45(6): 467-468, 1991; US Patent No. 4,432,921; EP 0 275 544; PCT/DK93/00217, 1992. Laccase is also useful in the copolymerization of lignin with low molecular weight compounds, such as is described in Appl. Microbiol. Biotechnol. 40: 760-767.

10 The laccase of the present invention can also be used for in-situ depolymerization of lignin in Kraft pulp, thereby producing a pulp with lower lignin content. This use of laccase is an improvement over the current use of chlorine for depolymerization of lignin, which leads to the production of chlorinated aromatic compounds, which are an environmentally undesirable by-product of paper mills. Such uses are described in, for example, Current opinion in Biotechnology 3: 261-266, 1992; J. Biotechnol. 25: 333-339, 1992; Hiroi et al., Svensk papperstidning 5: 162-166, 1976.

20 Since the environment in a paper mill is typically alkaline, the present laccase is more useful for this purpose than other known laccases, which function best under acidic conditions.

Oxidation of dyes or dye precursors and other

chromophoric compounds leads to decolorization of the compounds. Laccase can be used for this purpose, which can be particularly advantageous in a situation in which a dye transfer between fabrics is undesirable, e.g., in the textile industry and in the detergent industry. Methods for dye transfer inhibition and dye oxidation can be found in WO 92/01406; WO 92/18683; EP 0495836; Calvo, Mededelingen van de Faculteit Landbouw-wetenschappen/Rijiksuniversitet Gent. 56: 1565-1567, 1991; Tsujino et al., J. Soc. Chem. 42: 273-282, 1991. Use of laccase in oxidation of dye precursors

for hair dyeing is disclosed in U.S. Patent No. 3,251,742, the contents of which are incorporated herein by reference.

The present laccase can also be used for the polymerization or oxidation of phenolic compounds present in liquids. An example of such utility is the treatment of juices, such as apple juice, so that the laccase will accelerate a precipitation of the phenolic compounds present in the juice, thereby producing a more stable juice. Such applications have been described in Stutz, Fruit processing 7/93, 248-252, 1993; Maier et al., Dt. Lebensmittel-rindschau 86(5): 137-142, 1990; Dietrich et al., Fluss. Obst 57(2): 67-73, 1990.

Laccases such as the *Scytalidium* laccase are also useful in soil detoxification (Nannipieri et al., J.

15 Environ. Qual. <u>20</u>: 510-517,1991; Dec and Bollag, Arch. Environ. Contam. Toxicol. <u>19</u>: 543-550, 1990).

The invention is further illustrated by the following non-limiting examples.

EXAMPLES

20

I. ISOLATION OF SCYTALIDIUM THERMOPHILUM LACCASE GENE

A. MATERIALS AND METHODS

1. DNA Extraction and Hybridization analysis

Total cellular DNA is extracted from fungal cells of Scytalidium thermophila strain E421 grown 24 hours in 25 ml of YEG medium (0.5% yeast extract, 2% glucose) using the following protocol: Mycelia are collected by filtration through Miracloth (Calbiochem) and washed once with 25 ml of TE buffer. Excess buffer is drained from the mycelia which are subsequently frozen in liquid nitrogen. Frozen mycelia are ground to a fine powder in an electric coffee grinder, and the powder added to 20 ml of TE buffer and 5 ml of 20% SDS (w/v) in a disposable plastic centrifuge tube.

The mixture is gently inverted several times to ensure mixing, and extracted twice with an equal volume of phenol:chloroform:isoamyl alcohol (25:24:1). Sodium acetate (3M solution) is added to give a final concentration of 0.3 5 M and the nucleic acids are precipitated with 2.5 volumes of ice cold ethanol. The tubes are centrifuged at $15,000 \times g$ for 30 minutes and the pellet is allowed to air-dry for 30 minutes before resuspending in 0.5 ml of TE buffer. DNasefree ribonuclease A is added to a concentration of 100µg/ml 10 and the mixture is incubated at 37°C for 30 minutes. Proteinase K (200 μ g/ml) is added and each tube is incubated an additional one hour at 37°C. Finally, each sample is extracted twice with phenol:chloroform:isoamyl alcohol before precipitating the DNA with sodium acetate and 15 ethanol. DNA pellets are dried under vacuum, resuspended in TE buffer, and stored at 4°C.

Total cellular DNA samples are analyzed by Southern hybridization. Approximately 5µg of DNA is digested with EcoRI and fractionated by size on a 1% agarose gel. 20 is photographed under short wavelength UV and soaked for 15 minutes in 0.5 M NaOH, 1.5 M NaCl followed by 15 minutes in 1 M Tris-HCl, pH 8, 1.5 M NaCl. DNA in the gel is transferred onto Zeta-Probe™ hybridization membrane (BioRad Laboratories) by capillary blotting in 20 X SSPE (R. W. 25 Davis et al., Advanced Bacterial Genetics, A Manual for Genetic Engineering. Cold Spring Harbor Press. 1980) Membranes are baked for 2 hours at 80°C under vacuum and soaked for 2 hours in the following hybridization buffer at 45°C with gentle agitation: 5X SSPE, 35% formamide (v/v), 30~0.3% SCS, $200\mu g/ml$ denatured and sheared salmon testes DNA. The laccase-specific probe fragment (approx. 1.5 kb) encoding the 5'-portion of the N. crassa lcc1 gene is amplified from $N.\ crassa$ genomic DNA using standard PCR conditions (Perkin-Elmer Cetus, Emeryville, CA) with the

following pair of primers: forward primer, 5' CGAGACTGATAACTGGCTTGG 3'; reverse primer, 5' ACGGCGCATTGTCAGGGAAGT 3'. The amplified DNA segment is first cloned into a TA-cloning vector (Invitrogen, Inc., San 5 Diego, CA), then purified by agarose gel electrophoresis following digestion with EcoRI. The purified probe fragment is radiolabeled by nick translation with α[32P]dCTP(Amersham) and added to the hybridization buffer at an activity of approximately 1 X 106 cpm per ml of buffer. 10 incubated overnight at 45°C in a shaking water bath. Following incubation, the membranes are washed once in 0.2 X SSPE with 0.1% SDS at 45°C followed by two washes in 0.2 X SSPE(no SDS) at the same temperature. The membranes are allowed to dry on paper towels for 15 minutes, then wrapped 15 in Saran Wrap™ and exposed to x-ray film overnight at -70°C with intensifying screens (Kodak).

2. DNA Libraries and Identification of Laccase Clones Genomic DNA libraries are constructed in the bacteriophage cloning vector λ -EMBL4(J.A.Sorge, in Vectors, 20 A Survey of Molecular Cloning Vectors and Their Uses, Rodriguez et al., eds, pp.43-60, Butterworths, Boston, 1988). Briefly, total cellular DNA is partially digested with Sau3A and size-fractionated on low-melting point agarose gels. DNA fragments migrating between 9kb and 23 kb 25 are excised and eluted from the gel using ß-agarase (New England Biolabs, Beverly MA). The eluted DNA fragments are ligated with BamHI-cleaved and dephosphorylated λ -EMBL4 vector arms, and the ligation mixtures are packaged using commercial packaging extracts (Stratagene, LaJolla, CA). 30 The packaged DNA libraries are plated and amplified on Escherichia coli K802 cells. Approximately 10,000-20,000 plaques from each library are screened by plaquehybridization with the radiolabeled lcc1 DNA fragment using

the conditions described above. Plaques which give hybridization signals with the probe are purified twice on *E. coli* K802 cells, and DNA from the corresponding phage is purified from high titer lysates using a Qiagen Lambda 5 kit(Qiagen, Inc., Chatsworth, CA).

3. Analysis of Laccase Genes

Restriction mapping of laccase clones is done using standard methods (Lewin, Genes. 2d ed., Wiley & Sons, 1985, New York). DNA sequencing is done with an Applied

Biosystems Model 373A automated DNA Sequencer (Applied Biosystems, Inc., Foster City, CA) using the primer walking technique with dye-terminator chemistry (H. Giesecke et al., J. Virol. Methods 38: 47-60, 1992). Oligonucleotide sequencing primers are synthesized on an Applied Biosystems

model 394 DNA/RNA Synthesizer.

B. RESULTS AND DISCUSSION

1. Identification of Laccase Gene Sequence

Total cellular DNA samples are prepared from the

species Neurospora crassa, Botrytis cinerea, and
Scytalidium. Aliquots of these DNA preparations are
digested with BamHI and fractionated by agarose gel
electrophoresis. DNA in the gel is blotted to a Zeta-ProbeTM
membrane filter (BioRad Laboratories, Hercules,CA) and

probed under conditions of mild stringency with a
radiolabeled fragment encoding a portion of the N. crassa
lcc1 gene, as described above. Laccase-specific sequences
are detected in the genomes of S. thermophilum and the N.
crassa control, but not in the B. cinerea genomic DNA with
this probe.

2. Cloning and Characterization of Scytalidium thermophila Laccase (StL) Gene

The S. thermophilum laccase gene is isolated using plaque hybridization to screen the genomic DNA library made

in λ -EMBL4. The library contains approximately 250,000 independent clones before amplification, and 12,000 plaques are screened by hybridization with a radiolabeled N. crassa laccase gene fragment as described above. Nine plaques are 5 identified which hybridize strongly to the probe. DNA is isolated from four of these clones and analyzed by restriction mapping. All four contain a 3kb BamHI fragment that is originally identified in southern blotting with genomic DNA as described above. This fragment is isolated 10 from one clone and inserted into a pBluescript vector(Stratagene Cloning Systems, La Jolla, CA). DNA sequence analysis indicates that only a portion of the gene is located on this segment. Consequently, a 6 kb XhoI fragment which contains the entire lccs gene is subcloned 15 into pBluescript to derive the plasmid pShTh6. A restriction map of the 6 kb insert in this plasmid is shown in Figure 3. The nucleic acid sequence is shown in Figure 1 and SEQ ID NO: 1. The deduced amino acid sequence of StL is obtained on the basis of amino acid sequence homology with 20 N. crassa laccase. StL shares approximately 58% amino acid sequence identity with NcL, and this sequence similarity is highest among those amino residues that are involved in the formation of the active site copper center. StL, like NcL appears to be synthesized as a preproenzyme (616 amino acids 25 with a 21 amino acid signal peptide and a propeptide of 24 amino acids). However, since the amino terminal sequence of the mature StL protein is not yet determined, the exact length of the propeptide is not certain. There are five potential sites for N-linked glycosylation in StL. 30 potential C-terminal processing signal with homology to N. crassa laccase also exists in StL (Asp-Ser-Gly-Leu*Lys564) which may result in the proteolytic removal of the last seven amino acids from the primary translation product.

The presence of four small introns (63, 58, 55 and 65 nucleotides) is determined by comparing the open reading frames within the coding region of *lccS* to the primary structure of NcL. Excluding these intervening sequences, the coding region contains 60.8% G+C. The base composition of *lccS* reflects a bias for codons ending in G or C.

II. EXPRESSION OF SCYTALIDIUM LACCASE IN ASPERGILLUS

10 A. MATERIALS AND METHODS

1. Bacterial and Fungal Host Strains

Escherichia coli JM101 (Messing et al., Nucl. Acids Res. 9:309-321, 1981) is used as a host for construction and routine propagation of laccase expression vectors in this study. Fungal hosts for laccase expression included the Aspergillus niger strain Bo-1, as well as a uridine-requiring (pyrG) mutant of the α-amylase-deficient Aspergillus oryzae strain HowB104.

2. Plasmids

Plasmid pSHTh5 is a pBluescript(Stratagene Cloning Systems, LaJolla, CA) derivative which contains a 6kb XhoI fragment of S. thermophilum DNA encoding StL. Plasmid pToC68(WO 91/17243) contains the A. oryzae TAKA-amylase promoter and A. niger glaA terminator, and pToC90(WO 91/17243) carries the A. nidulans amdS gene.

3. Construction of Laccase Expression Vectors

The construction strategy for the laccase expression vector pShTh15 is outlined in Figure 2. The promoter directing transcription of the laccase gene is obtained from the A. oryzae α-amylase (TAKA-amylase) gene (Christensen et al., supra), and terminator from the A. niger glaA (glucoamylase) terminator region. The expression vector is constructed as follows. A 60 basepair synthetic DNA linker,

5 TCGAGATGAAGCGCTTCTTCATTAATAGCCTTCTCGCAGGGCTCCTCAACTCAGGGGCC 3 '

- 3' CTACTTCGCGAAGAGTAATTATCGGAAGACGAAGAGCGTCCCGAGGAGTTGAGTCC 5'
- including the region from start codon to an ApaI site, is inserted into XhoI- and ApaI-digested pBluescriptSK(Stratagene, LaJolla, CA) to produce an intermediate termed pShTh11.5. This vector is digested with ApaI and Asp718 and ligated with a 662 base pair ApaI-Asp718 fragment encoding a portion of StL from pShTh5, generating a second intermediate called pShTh13.1. An XbaI site is introduced immediately downstream of the stop codon using pShTh5 as a template for a PCR reaction with the following primers:forward:
 5'GTCATGAACAATGACCT 3'; reverse:
- 5'AGAGAGTCTAGATTAAACAATCCGCCCAACTAC3'. The amplified fragment is digested with NsiI and XbaI and subcloned into pUC518 to created the intermediate called pShTh12.8. The pShTh12.8 vector is digested with EcoRI and Asp718 and ligated with a 700 base pair EcoRI-Asp718 fragment from pShTh13.1 to generate pShTh13.1 to generate pShTh13.2. An 800 base pair NsiI-Asp718 fragment containing the final portion of the laccase coding region is obtained from pShTh5 and inserted into NsiI- and Asp718-cleaved pShTh13.2 to give pShTh14. Lastly, the 2.2 kb laccase coding region in pShTh14 is removed by cleavage with XhoI and XbaI and inserted between the XhoI and XbaI sites of pToC68 to generate the expression vector pShTh15.

4. Transformation of Aspergillus host cells

Methods for co-transformation of Aspergillus strains are as described in Christensen et al., supra. For introduction of the laccase expression vectors into A. oryzae HowB 104 pyrG, equal amounts (approximately 5 µg each) of laccase expression vector and pPyrG, which harbors the cloned A. nidulans pyrG gene, are used. Protrophic(Pyr+) transformants are selected on Aspergillus minimal medium

(Rowlands and Turner, Mol. Gen. Genet. 126: 201-216, 1973), and the transformants are screened for the ability to produce laccase on minimal medium containing 1 mM 2,2'-azinobis(3-ethylbenzthiazolinesulfonic acid)[ABTS]. Cells which secrete active laccase oxidize the ABTS, producing a green halo surrounding the colony. A. niger Bo-1 protoplasts are cotransformed using equal amounts (approximately 5µg each) of laccase expression vector and pToC90 which contains the A. nidulans amds (acetamidase) gene (Hynes et al., Mol. Cell Biol. 3: 1430-1439, 1983. AmdS+ transformants are selected on Cove minimal medium (Cove, Biochim. Biophys. Acta 113: 51-56, 1966) with 1% glucose as the carbon source and acetamide as the sole nitrogen source and screened for laccase expression on Cove medium with 1 mM ABTS.

5. Analysis of Laccase-Producing Transformants

Transformants which produce laccase activity on agar plates are purified twice through conidiospores and spore suspensions in sterile 0.01% Tween-80 are made from each. density of spores in each suspension is estimated 20 spectrophotometrically (A₅₉₅ nm). Approximately 0.5 absorbance units of spores are used to inoculate 25 ml of ASPO4 or MY50 medium in 125 ml plastic flasks. The cultures are incubated at 37°C with vigorous aeration (approximately 200 rpm) for four to five days. Culture broths are harvested by 25 centrifugation and the amount of laccase activity in the supernatant is determined using syringaldazine as a substrate. Briefly, 800 μ l of assay buffer (25 mM sodium acetate, pH 5.5, 40 μ M CuSo₄) is mixed with 20 μ l of culture supernatant and 60 μl of 0.28 mM syringaldazine stock solution (Sigma Chemical 30 Co., St. Louis, MO) in 50% ethanol. The absorbance at 530 nm is measured over time in a Genesys 5 UV-vis spectrophotometer (Milton-Roy). One laccase unit(LACU) is defined as the amount of enzyme which oxidizes one µmole of substrate per minute at room temperature. SDS-polyacrylamide gel

electrophoresis (PAGE) is done using precast 10-27% gradient gels from Novex (San Diego, CA). Protein bands are developed using Coomassie Brilliant Blue (Sigma).

5 B.RESULTS AND DISCUSSION

1. Expression of Scytalidium laccase

The expression vector pShTh15 is used in conjunction with pPyrG (A. nidulans pyrG) or pToC90(A. nidulans amdS) plasmids to generate A. oryzae and A. niger co-transformants which

10 express StL. As shown in Table 1, the number of laccase-producing co-transformants obtained in A. oryzae HowB104pyrG is small (3.7% of Pyr+ transformants) compared to the number obtained in A. niger Bo-1 using amdS selection (71.5% of AmdS+transformants). It is unknown whether this is due to an abnormally low co-transformation(i.e., integration) frequency or extremely low expression or laccase degradation in many A. oryzae transformants. Expression levels of StL range from about 50mg/l in shake flasks and 1-2g/l in a fermentor.

20

III. PURIFICATION AND CHARACTERIZATION OF RECOMBINANT SCYTALIDIUM LACCASE

A. MATERIALS AND METHODS

1. Materials

25 Chemicals used as buffers and substrates are commercial products of at least reagent grade. Chromatography is performed on either a Pharmacia FPLC. Spectroscopic assays are conducted on either a spectrophotometer(Shimadzu PC160) or a microplate reader(Molecular Devices). Britton & Robinson(B&R) buffers are prepared according to the protocol described in Quelle, Biochemisches Taschenbuch, H.M. Raven, II. Teil, S.93 u. 102, 1964.

2. Fermentation

A 1 ml aliquot of a spore suspension of Aspergillus oryzae transformant HowB104-pShTh15-2(approximately 109 spores/ml) is added aseptically to a 500 ml shake flask containing 100 ml of sterile shake flask medium (maltose, 5 50g/1; $MgSO_4 \cdot 7H_2O$, 2g/1; KH_2PO_4 , 10g/1; K_2SO_4 , 2g/1; $CaCl_2 \cdot 2H_2O$ 0.5 g/l; Citric acid, 2g/l; yeast extract, 10g/l; trace metals $[ZnSO_4 \cdot 7H_2O, 14.3 g/1; CuSO_4 \cdot 5H_2O, 2.5 g/1; NiCl_2 \cdot 6H_2O,$ 0.5 g/l; $FeSO_4 \cdot 7H_2O$, 13.8 g/l, $MnSO_4 \cdot H_2O$, 8.5 g/l; citric acid, 3.0 g/l], 0.5 ml/l; urea, 2g/l, made with tap water and 10 adjusted to pH 6.0 before autoclaving), and incubated at 37°C on a rotary shaker at 200 rpm for 18 hours. 50 ml of this culture is aseptically transferred to a 3 liter fermentor containing 1.8 liters of the fermentor media (MgSO₄·7H₂O, 2g/l; KH_2PO_4 , 2g/1; citric acid 4g/1; K_2SO_4 , 3g/1; $CaCl_2 \cdot 2H_2O$, 2g/1; 15 trace metals, 0.5 ml/l; pluronic antifoam, 1ml/l). fermentor temperature is maintained at 34°C by the circulation of cooling water through the fermentor jacket. Sterile air is sparged through the fermentor at a rate of 1.8 liter/min (1v/v/m). The agitation rate is maintained between 600 and 20 1300 rpm at approximately the minimum level required to maintain the dissolved oxygen level in the culture above 20%. Sterile feed (Nutriose 725[maltose syrup], 225 g/l; urea, 30 g/l; yeast extract, 15 g/l; pluronic antifoam, 1.5 ml/l, made up with distilled water and autoclaved) is added to the 25 fermentor by use of a peristaltic pump. The feed rate profile during the fermentation is as follows: 30 g of feed is added initially before inoculation; 0-24 h, 2 g/l h; 24-48 h, 4 g/l h; 48h-end, 6 g/1.

Copper(in the form of CuCl₂, CuSO4 or other soluble salt)
30 is made as a 400% stock in water or a suitable buffer, filter sterilized and added aseptically to the tank to a final level of 0.5 mM.

Samples for enzyme activity determination are withdrawn and filtered through Miracloth to remove mycelia. These samples are assayed for laccase activity by the LACU assay described above. Laccase activity is found to increase continuously during the course of the fermentation, with a value of approximately 3.6 LACU/ml achieved after 115 hours in the fermentation containing excess copper. At a specific activity of 1.9 LACU/mg, this corresponds to over 1.8 g/l recombinant laccase expressed by this transformant.

10 3. Enzymatic Assay

Laccase activity is determined by syringaldazine oxidation at 30°C in a 1-cm quartz cuvette. 60µl syringaldazine stock solution (0.28 mM in 50% ethanol) and 20 µl sample are mixed with 0.8 ml preheated buffer solution.

The oxidation is monitored at 530nm over 5 minutes. The activity is expressed as µmole substrate oxidized per minute. B&R buffers with various pHs are used. The activity unit is referred to here as "SOU". A buffer of 25 mM sodium acetate, 40 µM CuSO4, pH 5.5, is also used to determine the activity, which is referred to as LACU, as defined above. 2,2'-azinobis(3-ethylbenzo thiazoline-6-sulfonic acid) (ABTS) oxidation assays are done using 0.4 mM ABTS, B&R buffer, pH 4.1, at room temperature by monitoring AA405. An ABTS oxidase activity overlay assay is performed by pouring cooled ABTS-agarose(0.05 g ABTS, 1 g agarose, 50 ml H₂O, heated to

agarose (0.05 g ABTS, 1 g agarose, 50 ml H_2O , heated to dissolve agarose) over a native-IEF gel and incubating at room temperature. Thermostability analysis is performed using samples that have ~3 μ M enzyme preincubated for one hour in B&R buffer, at pH 2.7, 6.1, and 9.0, and various temperatures.

30 Samples are assayed after a 44-fold dilution into B & R buffer, pH 4.1, at room temperature.

3. Purification from a fermentor broth

1.2 liters of cheese-cloth filtered broth (pH 7.9, 13 mS) is filtered through Whatman #2 filter paper and concentrated

on a Spiral Concentrator(Amicon) with a S1Y100 membrane (MWCO:100) to 200 ml. The concentrate is adjusted to 0.86 mS by diluting it in water and reconcentrated on S1Y100 to 324 ml. The washed and concentrated broth has a dense greenish color.

The broth is frozen overnight at -20°C, thawed the next day(without any loss of activity) and loaded onto a Q-Sepharose XK26 column (120 ml), preequilibrated with 10 mM Tris, pH 7.7, 0.9 mS. The blue laccase band is eluted during a linear gradient with 2 M NaCl.

Pooled laccase fractions(44 ml), dialyzed in 3.5 liters of 10 mM NaAc, pH 5.5, 0.8 mS at 4°C overnight, are loaded onto a Mono-Q 16/10 (40 ml), preequilibrated with 10 mM MES, pH 5.3, 0.8 mS. The laccase eluted during a linear gradient with 1 M NaCl shows apparent homogeneity on SDS-PAGE.

4. Analysis of amino acid content and N-terminus

N-terminal sequencing is performed on an ABI 476A sequencer; and total amino acid analysis, from which the extinction coefficient of laccase is determined, is performed on a HP AminoQuant instrument.

B. RESULTS AND DISCUSSION

1. Purification

From 1200 ml fermentor broth, about 0.6g of laccase are isolated. Initial concentration using a membrane with MWCO of 100 kDa removes significant amounts of brown material and small contaminant proteins. The low affinity of the laccase toward Q-Sepharose matrix equilibrated with 10 mM Tris, pH 7.7, facilitates its separation from other impurities. The enriched fractions are further purified by Mono-Q at pH 5.3. Although it has a pI of 5.1, the laccase migrates slowly on Mono-Q and is separated from impurities during the washing by 10 mM MES, pH 5.3. An overall 15-fold purification and a recovery of 60% are achieved.

2. Characterization

The purified laccase shows a MW of 75-80 kDa on SDS-PAGE.

The difference between the MW derived from DNA sequence(63 kDa) and the observed MW is attributable to glycosylation.

5 Native IEF shows 3 bands near pI of about 5.1, which are active in ABTS overlay assay.

3. N-terminal sequencing

Directly sequencing the N-terminus of the purified laccase from samples either in desalted solution or on PVDF membrane are unsuccessful. This result suggests a blocked N-terminus, likely a pyroglutamate site based on the gene sequence.

The spectrum of the blue laccase has absorption maxima at 276 and 602 nm; with $Abs_{280}/Abs_{600}=23$ and $Abs_{330}/Abs_{589}=2.1$. The extinction coefficient determined by amino acid analysis is 1.9 1/(g*cm).

The activity is tested by using either syringaldazine or ABTS as substrates. Expressed as per Abs₂₈₀ or per mg, the laccase has a value of 2.2 or 4.2 units for SOU at pH 7, 20 respectively.

The pH profiles of laccase activity has optimal pH of 7 and 4, for syringaldazine and ABTS oxidation, respectively(Figure 4). Thermostability analysis at three pHs is shown in Figure 5. The laccase is more stable at neutral to alkaline pH than at acidic pH. Thermoactivation is also observed in neutral-alkaline pH range.

Deposit of Biological Materials

The following biological material has been deposited under the terms of the Budapest Treaty with the Agricultural Research Service Patent Culture Collection, Northern Regional Research Center, 1815 University Street, Peoria, Illinois, 61604 and given the following accession number.

Deposit

Accession Number

E. coli JM101 containing

pShTh15 NRRL B-21262

5

PCT/US95/06816 WO 95/33837

SEQUENCE LISTING

(1) GENERAL INFORMATION:

(i) APPLICANT:

- (A) NAME: Novo Nordisk Biotech, Inc.
- (B) STREET: 1445 Drew Avenue
- (C) CITY: Davis, California
- (D) COUNTRY: United States of America
- (E) POSTAL CODE (ZIP;: 95616-4880 (F) TELEPHONE: (916) 757-8100
- (G) TELEFAX: (916) 758-0317

(ii) TITLE OF INVENTION: PURIFIED SCYTALIDIUM LACCASES AND NUCLEIC ACIDS ENCODING SAME

(iii) NUMBER OF SEQUENCES: 2

- (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Novo Nordisk of North America, Inc.
 - (B) STREET: 405 Lexington Avenue, Suite 6400
 - (C) CITY: New York
 - (D) STATE: New York
 - (E) COUNTRY: U.S.A.
 - (F) ZIP: 10174-6401

(v) COMPUTER READABLE FORM:

- (A) MEDIUM TYPE: Floppy disk
- (B) COMPUTER: IBM PC compatible
- (C) OPERATING SYSTEM: PC-DOS/MS-DOS
- (D) SOFTWARE: PatentIn Release #1.0, Version #1.25 (EPO)

(vi) CURRENT APPLICATION DATA:

- (A) APPLICATION NUMBER: US
- (B) FILING DATE: 31-May-95
- (C) CLASSIFICATION:

(vii) PRIOR APPLICATION DATA:

- (A) APPLICATION NUMBER: US 08/253,784
- (B) FILING DATE: 03-June-1994

(viii) ATTORNEY/AGENT INFORMATION:

- (A) NAME: Lowney, Karen A.
 (B) REGISTRATION NUMBER: 31,274
 (C) REFERENCE/DOCKET NUMBER: 4186.204-WO

(ix) TELECOMMUNICATION INFORMATION:

- (A) TELEPHONE: 212 867 0123
- (B) TELEFAX: 212 867 0298

(2) INFORMATION FOR SEQ ID NO: 1:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2476 base pairs

 - (B) TYPE: nucleic acid (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Scytalidium thermophilum

(ix) FEATURE:

(A) NAME/KEY: intron

(B) LOCATION: 349..411

	(ix)	(A	TURE) NA) LO	ME/K	EY: ON:	intr 502.	on .559									
	(ix)	(A		ME/K	EY:											
	(ix)	(A	TURE) NA) LO	ME/K	EY:	intr 1739	on 18	04								
	(ix)	(A	TURE) NA) LO	ME/K	EY:	CDS join	(10 18	63 05	48, 2194	412.	.501	, 56	506	31,	6871738,	
	(xi)	SEQ	UENC	E DE	SCRI	PTIC	N: S	EQ I	D NC	: 1:						
CTGA	ATTI	'AA A	TACA	GGAA	G AI	CGCA	TTCA	ATC	CAGO	CTA	GACI	GCAC	AA T	GGT	CTGCA	60
CGAC	CGTC	GC A	CACC	TGCC	TA A	'AGTG	TTAA	TAA	CGGN	ICTA	ATAC	C AT Me	G Alet Ly	AG CC	GC TTC rg Phe	117
TTC Phe 5	ATT Ile	AAT Asn	AGC Ser	CTT Leu	CTG Leu 10	CTT Leu	CTC Leu	GCA Ala	GGG Gly	CTC Leu 15	CTC Leu	AAC Asn	TCA Ser	GGG Gly	GCC Ala 20	165
CTC Leu	GCG Ala	GCT Ala	CCG Pro	TCT Ser 25	ACA Thr	CAT His	CCC Pro	AGA Arg	TCA Ser 30	AAC Asn	CCC Pro	GAC Asp	ATA Ile	CTG Leu 35	CTT Leu	213
GAA Glu	AGA Arg	GAT Asp	GAC Asp 40	CAC His	TCC Ser	CTT Leu	ACG Thr	TCT Ser 45	CGG Arg	CAA Gln	GGT Gly	AGC Ser	TGT Cys 50	CAT His	TCT Ser	261
CCA Pro	AGC Ser	AAC Asn 55	CGC Arg	GCC Ala	TGT Cys	TGG Trp	TGC Cys 60	TCT Ser	GGC Gly	TTC Phe	GAT Asp	ATC Ile 65	AAC Asn	ACG Thr	GAT Asp	309
TAT Tyr	GAG Glu 70	ACC Thr	AAG Lys	ACT Thr	CCA Pro	AAC Asn 75	ACC Thr	GGA Gly	GTG Val	GTG Val	CGG Arg 80	CGG Arg	GTT	AGTA:	rcc	358
CAAC	TTAC	GT I	TGAC	CAAC	SA AZ	\TGG/	ACGTO	AAC	TGT	CTG	ACTO	CTCC	CGC !	rag		411
TAC Tyr	ACC Thr	TTT Phe	GAT Asp 85	ATC Ile	ACC Thr	GAA Glu	GTC Val	GAC Asp 90	AAC Asn	CGC	CCC Pro	GGT Gly	CCC Pro 95	GAT Asp	GGG	459
GTC Val	ATC Ile	AAG Lys 100	GAG Glu	AAG Lys	CTC Leu	ATG Met	CTT Leu 105	ATC Ile	AAC Asn	GAC Asp	AAA Lys	CTC Leu 110	CTG Leu	GTA	3G .	506
GTC	CTCTC	CGA A	ACGC	CTGC	er c	rgcc:	ACAC	A GC	GTAA	AACT	AAC	GAAC	cgc '	TAG		559
GGC Gly	CCG Pro	ACA Thr	GTC Val 115	TTC Phe	GCA Ala	AAC Asn	TGG Trp	GGC Gly 120	GAC Asp	ACC Thr	ATC Ile	GAG Glu	GTG Val 125	ACC Thr	GTC Val	607
AAC Asn	AAC Asn	CAC His 130	CTG Leu	AGA Arg	ACC Thr	AAC Asn	GGA Gly 135	GTA	AGCG'	TTC (GGAC:	ACAA	ag c	CCAG	CAACC	661

TAG	ACAC	ACT ·	CAAC	TGAC	CA A	GTAG									CAC His	 716
													GAG Glu			764
													CGC Arg 175			812
													GGC Gly			860
													CCC Pro			908
ATC Ile 210	GAC Asp	CTC	GGC Gly	GTC Val	CTC Leu 215	CCG Pro	CTG Leu	CAG Xaa	GAC Asp	TGG Trp 220	TAC Tyr	TAC Tyr	AAG Lys	TCC Ser	GCC Ala 225	956
													CCG Pro			1004
													ACT Thr 255			1052
													CAT His			1100
													TCG Ser			1148
													GTC Val			1196
													TAT Tyr			1244
													AAC Asn 335			1292
TTT Phe	GGA Gly	GGG Gly 340	CAG Gln	CAG Gln	AAG Lys	TGC Cys	GGC Gly 345	TTC Phe	TCG Ser	CAC His	AAT Asn	CCG Pro 350	GCG Ala	CCG Pro	GCA Ala	1340
													ACG Thr			1388
													GAT Asp			1436
													AAA Lys			1484

GGC Gly	AAT Asn	ACG Thr	CTG Leu 405	CCG Pro	GTG Val	ACG Thr	CTC Leu	CAT His 410	GTT Val	GAC Asp	CAG Gln	GCC Ala	GCG Ala 415	GCT Ala	CCA Pro	1532
CAC His	GTG Val	TTT Phe 420	ACG Thr	TGG Trp	AAG Lys	ATC Ile	AAC Asn 425	GGG Gly	AGC Ser	GCT Ala	GCG Ala	GAC Asp 430	GTG Val	GAC Asp	TGG Trp	1580
GAC Asp	AGG Arg 435	CCG Pro	GTG Val	CTG Leu	GAG Glu	TAT Tyr 440	GTC Val	ATG Met	AAC Asn	AAT Asn	GAC Asp 445	CTG Leu	TCT Ser	AGC Ser	ATT Ile	1628
CCG Pro 450	GTC Val	AAG Lys	AAC Asn	AAC Asn	ATT Ile 455	GTG Val	AGG Arg	GTG Val	GAC Asp	GGA Gly 460	GTC Val	AAC Asn	GAG Glu	TGG Trp	ACG Thr 465	1676
TAC Tyr	TGG Trp	CTC Leu	GTC Val	GAA Glu 470	AAC Asn	GAC Asp	CCG Pro	GAG Glu	GGC Gly 475	CGC Arg	CTC Leu	AGT Ser	TTG Leu	CCG Pro 470	CAT His	1724
	ATG Met				GTA	AGTC	ACA :	rccc	CCAC!	ra co	CATT	CGGA	A TG	ACCA	CCAG	1779
GTA	CTGA	CAC (CCTC	CTCC'	rc Ai	ATAG	GGA Gly	CAC His	GAT Asp	TTC Phe 480	TTT Phe	GTC Val	CTA Leu	GGC Gly	CGC Arg 485	1831
TCC Ser	CCC Pro	GAC Asp	GTC Val	TCG Ser 490	CCC Pro	GAT Asp	TCA Ser	GAA Glu	ACC Thr 495	CGC Arg	TTC Phe	GTC Val	TTT Phe	GAC Asp 500	CCG Pro	1879
GCC Ala	GTC Val	GAC Asp	CTC Leu 505	CCC Pro	CGT Arg	CTG Leu	CGC Arg	GGA Gly 510	CAC His	AAC Asn	CCC Pro	GTC Val	CGG Arg 515	CGC Arg	GAC Asp	, 1927
GTC Val	ACC Thr	ATG Met 520	CTT Leu	CCC Pro	GCG Ala	CGC Arg	GGC Glu 525	TGG Trp	CTG Leu	CTG Leu	CTG Leu	GCC Ala 530	TTC Phe	CGC Arg	ACG Thr	1975
GAC Asp	AAC Asn 535	Pro	GGC Gly	GCG Ala	TGG Trp	TTG Leu 540	TTC Phe	CAC His	TGC Cys	CAC His	ATC Ile 545	GCG Ala	TGR Trp	CAC His	GTG Val	·2023
TCG Ser 550	Gly	GGG Gly	TTA Leu	AGC Ser	GTC Val 555	Asp	TTT Phe	CTG Leu	GAG Glu	CGG Arg 560	CCG Pro	GAC Asp	GAG Glu	CTG Leu	CGC Arg 565	2071
GGG Gly	CAG Gln	CTG Leu	ACG Thr	GGA Gly 570	GAG Glu	AGC Ser	AAG Lys	GCG Ala	GAG Glu 575	TTG Leu	GAG Glu	CGT Arg	GTT Val	TGT Cys 580	CGC Arg	2119
GAG Glu	TGG Trp	AAG Lys	GAT Asp 585	Trp	GAG Glu	GCG Ala	AAG Lys	AGC Ser 590	Pro	CAT His	GGG	AAG Lys	ATC Ile 595	GAT Asp	TCG Ser	2167
GGG Gly	TTG	AAG Lys 600	Gln	CGG Arg	CGA Arg	TGG	GAT Asp 605	Ala	TGA	ggta	GTT	GGGC	GGAT	TG		2214
TTT	AACA	CGT	AGTG	GGTA	AG G	TTGG	GGCG	G GI	TTGT	TTGG	CGT	TTTC	AGG	GGTT	GGGGTG	2274
CGG	ATGC	TGG	TCAT	CCGG	GA A	ACGG	CTCT	A CA	ACTG	GTGT	CAA	TAGA	CTA	ATAT	AGAGTG	2334
ATC	AAAG	AAC	TGAG	GTTC	TG A	AAGA	GGCG	T GG	AAGT	CGCG	TTG	TGAC	TCC	CTTT	GCCATG	2394
TTG	GGAA	.GTG	TGGC	TCAA	CA I	TGTG	TTCA	G GI	TTGC	TCAG	GGI	GATN	TCG	AACT	GACGTN	2454

2476

PCT/US95/06816 WO 95/33837

TTGATGAGGG TTATTGCNTA GA

- (2) INFORMATION FOR SEQ ID NO: 2:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 616 amino acids (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Scytalidium thermophilum
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

Met Lys Arg Phe Phe Ile Asn Ser Leu Leu Leu Ala Gly Leu Leu

Asn Ser Gly Ala Leu Ala Ala Pro Ser Thr His Pro Arg Ser Asn Pro

Asp Ile Leu Leu Glu Arg Asp Asp His Ser Leu Thr Ser Arg Gln Gly

Ser Cys His Ser Pro Ser Asn Arg Ala Cys Trp Cys Ser Gly Phe Asp 50 55 60

Ile Asn Thr Asp Tyr Glu Thr Lys Thr Pro Asn Thr Gly Val Val Arg

Arg Tyr Thr Phe Asp Ile Thr Glu Val Asp Asn Arg Pro Gly Pro Asp

Gly Val Ile Lys Glu Lys Leu Met Leu Ile Asn Asp Lys Leu Leu Gly

Pro Thr Val Phe Ala Asn Trp Gly Asp Thr Ile Glu Val Thr Val Asn

Asn His Leu Arg Thr Asn Gly Thr Ser Ile His Trp His Gly Leu His

Gln Lys Gly Thr Asn Tyr His Asp Gly Ala Asn Gly Val Thr Glu Cys

Pro Ile Pro Pro Gly Gly Ser Arg Val Tyr Ser Phe Arg Ala Arg Gln

Tyr Gly Thr Ser Trp Tyr His Ser His Phe Ser Ala Gln Tyr Gly Asn 185

Gly Val Ser Gly Ala Ile Gln Ile Asn Gly Pro Ala Ser Leu Pro Tyr

Asp Ile Asp Leu Gly Val Leu Pro Leu Gln Asp Trp Tyr Tyr Lys Ser

Ala Asp Gln Leu Val Ile Glu Thr Leu Ala Lys Gly Asn Ala Pro Phe

Ser Asp Asn Val Leu Ile Asn Gly Thr Ala Lys His Pro Thr Thr Gly

Glu Gly Glu Tyr Ala Ile Val Lys Leu Thr Pro Asp Lys Arg His Arg

Leu Arg Leu Ile Asn Met Ser Val Glu Asn His Phe Gln Val Ser Leu 285 Ala Lys His Thr Met Thr Val Ile Ala Ala Asp Met Val Pro Val Asn 295 Ala Met Thr Val Asp Ser Leu Phe Met Ala Xaa Gly Gln Arg Tyr Asp 310 Val Thr Ile Asp Ala Ser Gln Ala Val Gly Asn Tyr Trp Phe Asn Ile Thr Phe Gly Gly Gln Gln Lys Cys Gly Phe Ser His Asn Pro Ala Pro Ala Ala Ile Phe Arg Tyr Glu Gly Ala Pro Asp Ala Leu Pro Thr Asp Pro Gly Ala Ala Pro Lys Asp His Gln Cys Leu Asp Thr Leu Asp Leu Ser Pro Val Val Gln Lys Asn Val Pro Val Asp Gly Phe Val Lys Glu Pro Gly Asn Thr Leu Pro Val Thr Leu His Val Asp Gln Ala Ala Ala Pro His Val Phe Thr Trp Lys Ile Asn Gly Ser Ala Ala Asp Val Asp Trp Asp Arg Pro Val Leu Glu Tyr Val Met Asn Asn Asp Leu Ser Ser Ile Pro Val Lys Asn Asn Ile Val Arg Val Asp Gly Val Asn Glu Trp Thr Tyr Trp Leu Val Glu Asn Asp Pro Glu Gly Arg Leu Ser Leu Pro His Pro Met His Leu His Gly His Asp Phe Phe Val Leu Gly Arg Ser Pro Asp Val Ser Pro Asp Ser Glu Thr Arg Phe Val Phe Asp Pro Ala Val Asp Leu Pro Arg Leu Arg Gly His Asn Pro Val Arg Arg Asp Val Thr Met Leu Pro Ala Arg Gly Trp Leu Leu Leu Ala Phe Arg Thr Asp Asn Pro Gly Ala Trp Leu Phe His Cys His Ile Ala Trp His Val Ser Gly Gly Leu Ser Val Asp Phe Leu Glu Arg Pro Asp Glu Leu Arg Gly Gln Leu Thr Gly Glu Ser Lys Ala Glu Leu Glu Arg Val Cys Arg Glu Trp Lys Asp Trp Glu Ala Lys Ser Pro His Gly Lys Ile Asp Ser Gly 600 Leu Lys Gln Arg Arg Trp Asp Ala

Applicant's or agent's file reference number	4186.204-WO	International application N TBA	

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13 bis)

A. The indications made below relate to the microorganism referm on page	ed to in the description
B. IDENTIFICATION OF	Further deposits are identified on an additional sheet
Name of depository institution Agricultural Research Service Patent Culture	Collection (NRRL)
Address of depository institution (including postal code and coun	try)
Northern Regional Research Center 1815 University Street Peoria, IL 61604, US	,
Date of deposit May 25, 1995	Accession Number NRRL B-21262
C. ADDITIONAL INDICATIONS (leave blank if not applications)	ble) This information is continued on an additional sheet
In respect of those designations in which a E during the pendency of the patent application only to be provided to an independent expert (Rule 28(4) EPC/Regulation 3.25 of Australia	, a sample of the deposited microorganism is nominated by the person requesting the sample
D. DESIGNATED STATES FOR WHICH INDICATIONS A	RE MADE (if the indications are not for all designated States)
E. SEPARATE FURNISHING OF INDICATIONS (leave blan	nk if not applicable)
The indication listed below will be submitted to the International "Accession Number of Deposit")	Bureau Later (specify the general nature of the indications e.g.
For receiving Office use only	For International Bureau use only
This sheet was received with the international application	☐ This sheet was received with the International Bureau on:
Authorized officer Dorls L. Brock Aus PCT International Division	Authorized officer
Form PCT/RO/134 (July 1992)	

35/1

WO 95/33837 PCT/US95/06816

What we claim is:

1. A DNA construct containing a sequence encoding a Scytalidium laccase.

- 2. The construct of Claim 1 which comprises a sequence encoding a Scytalidium thermophilum laccase.
- 3. The construct of Claim 1 which comprises a sequence encoding the amino acid sequence depicted in SEQ ID NO. 2.
 - 4. The construct of Claim 1, which comprises the nucleic acid sequence depicted in SEQ ID NO. 1.
- 15 5. The construct of Claim 1, which comprises the nucleic acid sequence contained in NRRL B-21262.
 - 6. A substantially pure Scytalidium laccase enzyme.
- 7. The enzyme of Claim 6 which is a *Scytalidium thermophilum* laccase.
- 8. The enzyme of Claim 6 which comprises the sequence depicted in SEQ ID NO. 2, or a sequence with at least about 80% homology thereto.
 - 9. A recombinant vector comprising a DNA construct containing a sequence encoding a Scytalidium laccase.
- 10. The vector of Claim 9 in which the sequence is operably linked to a promoter sequence.
 - 11. The vector of Claim 10 in which the promoter is a fungal or yeast promoter.

WO 95/33837 PCT/US95/06816

12. The vector of Claim 11 in which the promoter is the TAKA amylase promoter of Aspergillus oryzae.

- 5 13. The vector of Claim 11 in which the promoter is the glucoamylase (glaA;) promoter of Aspergillus niger or Aspergillus awamori.
- 14. The vector of Claim 9 which also comprises a selectable 10 marker.
 - 15. The vector of Claim 14 in which the selectable marker is selected from the group consisting of amdS, pyrG, argB, niaD, sC, and hygB.
- 16. The vector of Claim 14 in which the selectable marker is the amdS marker of Aspergillus nidulans or Aspergillus oryzae, or the pyrG marker of Aspergillus nidulans, Aspergillus niger, Aspergillus awamori, or Aspergillus oryzae.

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- 17. The vector of Claim 14 which comprises both the TAKA amylase promoter of Aspergillus oryzae and the amdS or pyrG marker of Aspergillus nidulans or Aspergillus oryzae.
- 25 18. A recombinant host cell comprising a heterologous DNA construct containing a nucleic acid sequence encoding a Scytalidium laccase.
 - 19. The host cell of Claim 18 which is a fungal cell.
 - 20. The host cell of Claim 19 which is an Aspergillus cell.
 - 21. The host cell of Claim 18 in which the construct is integrated into the host cell genome.

WO 95/33837 PCT/US95/06816

22. The host cell of Claim 18 in which the construct is contained on a vector.

- 5 23. The host cell of Claim 18 which comprises a construct containing a sequence encoding the amino acid sequence depicted in SEQ ID NO. 2.
- 24. A method for obtaining a laccase enzyme which comprises

 10 culturing a host cell comprising a DNA construct containing a
 sequence encoding a Scytalidium laccase enzyme, under
 conditions conducive to expression of the enzyme, and
 recovering the enzyme from the culture.
- 15 25. A Scytalidium enzyme obtained by the method of Claim 24.
 - 26. A method for polymerizing a lignin or lignosulfate substrate in solution which comprises contacting the substrate with a *Scytalidium* laccase.

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- 27. A method for in situ depolymerization in Kraft pulp which comprises contacting the pulp with a *Scytalidium* laccase.
- 28. A method for oxidizing dyes or dye precursors which 25 comprises contacting the dye or dye precursor with a Scytalidium laccase.
- 29. A method of polymerizing or oxidizing a phenolic compound which comprises contacting the phenolic compound with a scytalidium laccase.

1	CTGAATTTAAATACAGGAAGATCGCATTCAATCCAGCCTAGACTGCACAATGGTTCTGCA													60 1			
61 1	α	GACCO	GTCG(CACAC	CCTGC	CAAT	FAGTO	ATT:	ATAA(CGCC	TAAT	FACC	ATG M	AAG K	CGC R	TT F	116 4
117 4	С	TTC F	ATT I	AAT N	AGC S	CTT L	CTG L	CTT L	CTC L	GCA A	GGG G	CTC L	CTC L	AAC N	TCA S	GG G	161 19
162 19	G	GCC A	CTC L	GCG A		CCG P	TCT S	ACA T	CAT H	CCC P	AGA R	TCA S	AAC N	CCC P	GAC D	AT I	206 34
207 34	A	CTG L	CTT L		AGA R	GAT D	GAC D	CAC H	TCC S	CTT L	ACG T	TCT S	CGG R	CAA Q	GGT G	AG S	251 49
252 49	С	TGT C	CAT H	TCT S	CCA P	AGC S	AAC N	CGC R	GCC A	TGT C	TGG W	TGC C	TCT S	GGC G	TTC F	GA D	296 64
297 64	T	ATC I	AAC N	ACG T	GAT D		GAG E	ACC T	AAG K	ACT T	CCA P	AAC N	ACC T	GGA G	GTG V	GT V	341 79
342 79	G	CGG R	CGG R	GTT	AGTA'	TCCC	AAGT"	TACG	TTTG	ACCA	AGAA	ATGG	ACGT(GAAG	TGTG(CTG	398 81
700																	
399 81	A(CTCT	CCCG	CTAG	TAC	ACC T	TTT F	GAT D	ATC I	ACC T	GAA E	GTC V	GAC D	AAC N	CGC R	CC P	446 12
	A(C				Y	T	_	D	I	T	E	٧	D	N	R	Р	_
81 447 12 492	C	GGT G	CCC P	GAT D CTG	Y GGG G	GTC V	F ATC	D AAG K	I GAG E	T AAG K	E CTC L	V ATG M	D CTT L	N ATC I	R AAC N	GA D	12 491
81 447 12 492 27	C	GGT G AAA K	CCC P CTC L	GAT D CTG L	Y GGG G	GTC V TAGG	F ATC I	D AAG K TCTC	GAG E GAAC	T AAG K GCCT(E CTC L GCGT(V ATG M CTGC	D CTT L CACA(ATC I CAGC	R AAC N GTAA	GA D AACT	12 491 27 547 31
81 447 12 492 27 548 31	C C	GGT G AAA K ACGA	CCC P CTC L ACCG	GAT D CTG L CTAG	GGG G G GC	T GTC V TAGG	F ATC I GTCC	AAG K TCTC GTC V	GAAC GAAC TTC (F	AAG K GCCT(CTC L GCGT(AAC N	ATG M CTGC TGG (W AC G	CTT L CACAG	ATC I CAGCO	R AAC N GTAAA ACC A	GA D AACT	12 491 27 547 31 595

FIG.1A

1/8 SUBSTITUTE SHEET (RULE 26)

TGG CAC GGC TTG CAC CAA AAA GGA ACC AAC TAC CAC GAC GGC GCC 743 19 Y H D G T N K Н G L Н Q AAC GGC GTG ACC GAG TGT CCC ATC CCG CCC GGT GGC TCC CGA GTC 788 Р G S 34 Ρ Ρ I Ε C. ٧ T 19 TAC AGC TTC CGA GCG CGC CAA TAT GGA ACG TCA TGG TAC CAC TCC 833 G Ţ S R 0 Υ F 34 CAC TTC TCC GCC CAG TAT GGC AAC GGC GTG AGC GGC GCC ATC CAG 878 64 N G · V S G Α I Υ G S ATC AAC GGA CCC GCC TCC CTG CCC TAC GAC ATC GAC CTC GGC GTC 923 79 G S L P Υ D I D G CTC CCG CTG CAG GAC TGG TAC TAC AAG TCC GCC GAC CAG CTC GTC 968 94 S Υ Υ Κ D W Р L 0 ATC GAG ACC CTG GCC AAG GGC AAC GCT CCG TTC AGC GAC AAC GTC 1013 Р F 109 G N K E T L I CTC ATC AAC GGC ACC GCA AAG CAC CCC ACC ACT GGC GAA GGG GAG 1058 124 E G T T G K Н Р N G T Α I TAC GCC ATC GTG AAG CTC ACC CCG GGC AAA CGC CAT CGC CTG CGG 1103 1059 139 Κ R Н L T Ρ G ٧ K 124 I CTC ATC AAC ATG TCG GTG GAG AAC CAC TTC CAG GTC TCG CTG GCG 1148 1104 F 0 ٧ 154 E N Н S S ٧ 139 N I AAG CAC ACC ATG ACG GTC ATC GCG GCG GAC ATG GTC CCC GTC AAC 1193 1149 169 M ٧ Ρ ٧ A D T ٧ I Α T M 154 Н GCC ATG ACC GTC GAC AGC CTG TTT ATG GCC GNC GGG CAG CGG TAT 1238 1194 R 184 X G Q Y A F 169 T ٧ D S L GAT GTT ACC ATC GAC GCG AGC CAG GCG GTG GGG AAT TAC TGG TTC 1283 1239 G W F 199 N Υ S 0 Α ٧ D Α T Ī 184 ٧ AAC ATC ACC TIT GGA GGG CAG CAG AAG TGC GGC TTC TCG CAC AAT 1328 1284 QQKC G F S H 214 F G G T 199 I

FIG.1B

2/8

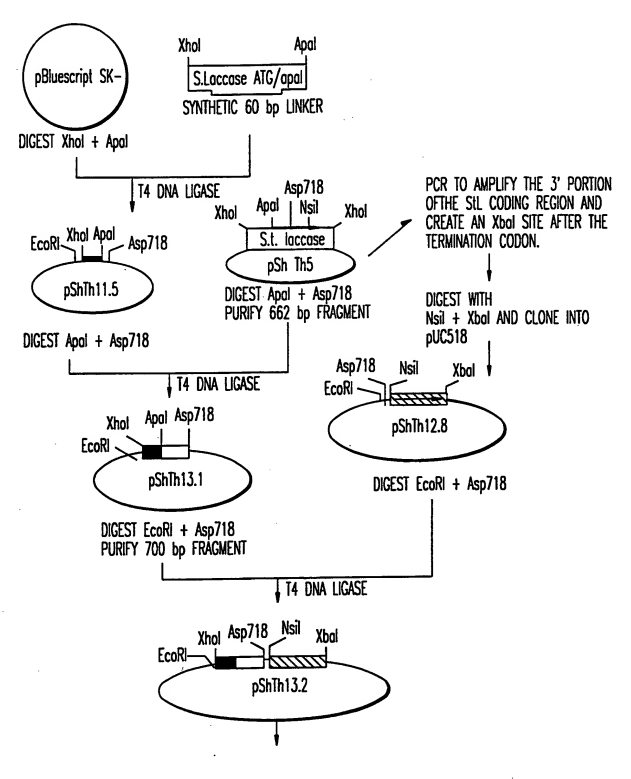
CCG GCG CCG GCA GCC ATC TTT CGC TAC GAG GGC GCT CCT GAC GCT 1373 1329 229 F R Υ Ε G Α D Р I 214 A Α CTG CCG ACG GAT CCT GGC GCT GCG CCA AAG GAT CAT CAG TGC CTG 1418 C 244 Α Р K D Н D Α GAC ACT TTG GAT CTT TCA CCG GTG GTG CAA AAG AAC GTG CCG GTT 1463 259 K N ٧ Р D S 244 D T 1464 GAC GGG TTC GTC AAA GAG CCT GGC AAT ACG CTG CCG GTG ACG CTC 1508 274 G Ε N 259 F CAT GTT GAC CAG GCC GCG GCT CCA CAC GTG TTT ACG TGG AAG ATC 1553 1509 T 289 Ρ Н 274 0 Α Α D Α AAC GGG AGC GCT GCG GAC GTG GAC TGG GAC AGG CCG GTG CTG GAG 1598 304 R D D 289 G S Α Α TAT GTC ATG AAC AAT GAC CTG TCT AGC ATT CCG GTC AAG AAC AAC 1643 1599 319 S S L 304 N N D ATT GTG AGG GTG GAC GGA GTC AAC GAG TGG ACG TAC TGG CTC GTC 1688 Υ 334 Ε Τ ٧ N 319 ٧ D G ٧ R GAA AAC GAC CCG GAG GGC CGC CTC AGT TTG CCG CAT CCG ATG CAT 1733 1689 349 PΕ G R L S 334 1734 CTA CAC GTAAGTCACATCCCCCACTACCATTCGGAATGACCACCAGGTACTGACACC 1790 351 349 L Н CTCCTCCTCAATAG GGA CAC GAT TTC TTT GTC CTA GGC CGC TCC CCC G 1838 12 F V. L G G Н F 351 1839 AC GTC TCG CCC GAT TCA GAA ACC CGC TTC GTC TTT GAC CCG GCC G 1883 Τ R F ٧ F D S Ε S Р ٧ TC GAC CTC CCC CGT CTG CGC GGA CAC AAC CCC GTC CGG CGC GAC G 1928 42 N R G Н R L 27 D L 1929 TC ACC ATG CTT CCC GCG CGC GGC TGG CTG CTG GCC TTC CGC A 1973 A R G W L L · L Α Р 42 V M L

FIG.1C

3/8
SUBSTITUTE SHEET (RULE 26)

1974	CG	GAC	AAC	CCG	GGC	GCG	TGG	TTG	TTC	CAC	TGC	CAC	ATC	GCG	TGR	С	2018
57	T	D	N	P	G	A	W	L	F	H	C	H	I	A	X		72
2019	AC	GTG	TCG	GGC	GGG	TTA	AGC	GTC	GAC	TTT	CTG	GAG	CGG	CCG	GAC	G	2063
72	H	V	S	G	G	L	S	V	D	F	L	E	R	P	D		87
2064	AG	CTG	CGC	GGG	CAG	CTG	ACG	GGA	GAG	AGC	AAG	GCG	GAG	TTG	GAG	С	2108
87	E	L	R	G	Q	L	T	G	E	S	K	A	E	L	E		102
2109	GT	GTT	TGT	CGC	GAG	TGG	AAG	GAT	TGG	GAG	GCG	AAG	AGC	CCG	CAT	G	2153
102	R	V	C	R	E	W	K	D	W	E	A	K	S	P	H		117
2154	GG	AAG	ATC	GAT	TCG	GGC	TTG	AAG	CAG	CGG	CGA	TGG	GAT	GCG	TGA	G	2198
117	G	K	I	D	S	G	L	K	Q	R	R	W	D	A	*		131
2199 131	GT	AGTT	GGGC	GGAT	TGTT	TAAC	ACGTA	AGTG(GTA	AGGT	TGGG(GCGG(STTTO	STTT	GCC 1	ſΤ	2258 131
2259 131	TT	CAGG	GGTT(GGCG.	TGCG(GATG(CTGG [*]	(CAT	CCGG(GAAA(CGGC	TCTA(CAAC	TGGT(STCA	AT .	2318 131
2319 131	AG	ACTA	ATATA	AGAG'	TGAT(CAAA(GAAC [*]	TGAG(STTC	[GAA/	AGAG(CCTC	GAA	GTCG(XTTX	;T	2378 131
2379 131	GA	CTCC	CTTT(GCCA [*]	TGTT(GGA	AGTG"	TGGC	(CAA	CATT(GTGT	TCAG(GTTT(GCTC/	\GGG]	[G	2438 131
2439 131	AT	NTCG	AACT(GACG [*]	TNT TO	GATG	AGGG	TAT.	IGC.	• • • •							2471 131

FIG.1D



5/8 FIG. 2A SUBSTITUTE SHEET (RULE 26)

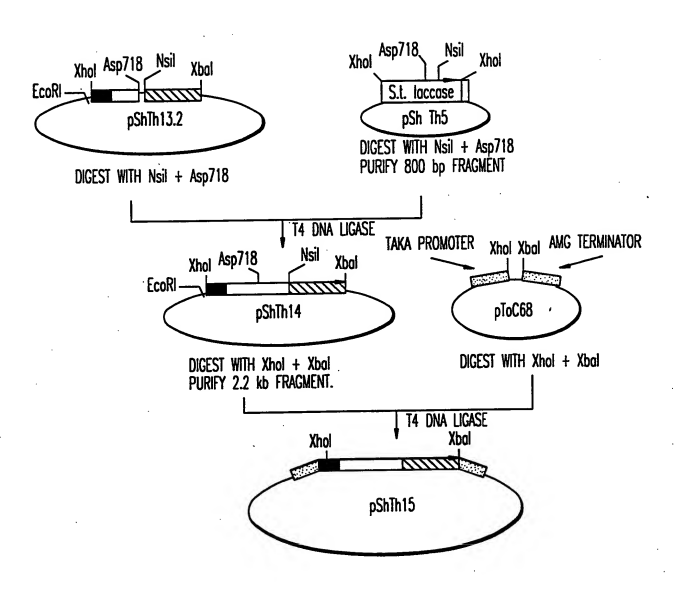
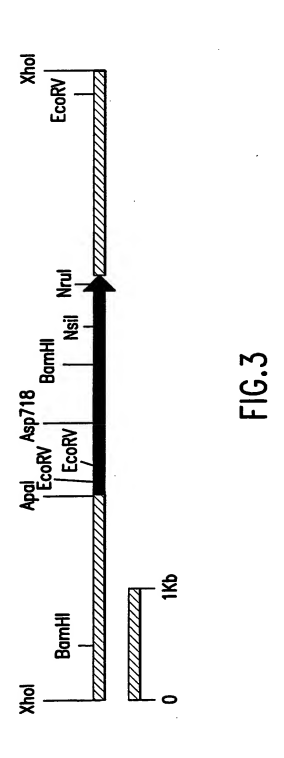
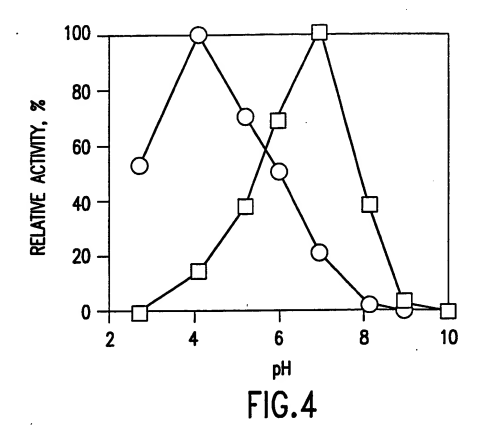


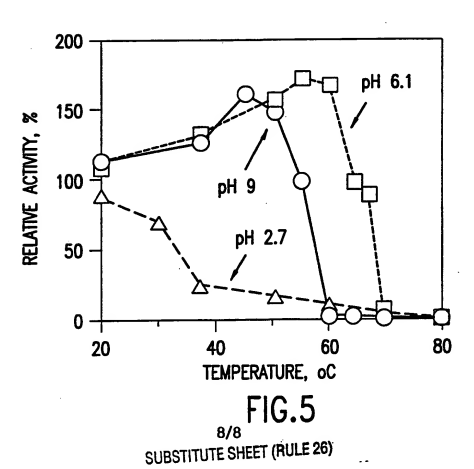
FIG.2B

6/8
SUBSTITUTE SHEET (RULE 26)



7/8 SUBSTITUTE SHEET (RULE 26)





A. CLASSIFICATION OF SUBJECT MATTER IPC 6 C12N15/53 C12N9/02 A61K7/13 C12N1/38 C12N1/15 //(C12N1/15,C12R1:66) A61K7/06 D21C5/00 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) C12N A61K D21C IPC 6 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Citation of document, with indication, where appropriate, of the relevant passages Category * 1,2,6,7, P,O, X ABSTRACTS OF PAPERS, 9,10, vol.209, no.1-2, April 1995 BERKA R. ET AL. 'Cloning of laccases from 18-20, 24,25 the thermophilic fungi Myceliophthora thermophila and Scytalidium thermophilum and their heterlogous expression in Aspergillus oryzae' see BIOT 196 JOURNAL OF BIOLOGICAL CHEMISTRY, A vol.263, no.2, 1988, BALTIMORE, MD US pages 885 - 896
GERMANN U. ET AL. 'Characterization of two allelic forms of Neurospora crassa laccase' see the whole document Patent family members are listed in annex. Further documents are listed in the continuation of box C. "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the Special categories of cited documents: "A" document defining the general state of the art which is not considered to be of particular relevance invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to "E" earlier document but published on or after the international filing date involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docucitation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled other means document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of mailing of the international search report Date of the actual completion of the international search 0 3. 10. 95 **29 August 1995** Authorized officer Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Espen, J Fax: (+31-70) 340-3016